Day: Friday Date: 2/23/2007

Time: 18:07:37

PALM INTRANET

Inventor Information for 10/809192

Inventor Name	City	State/Country
REDDY, MANNE SATYANARAYANA	HYDERABAD	INDIA
RAJAN, SRINIVASAN THIRUMALAI	HYDERABAD	INDIA
RAO, UPPALA VENKATA BHASKARA	HYDERABAD	INDIA
REDDY, KONDA SRINIVASA	HYDERABAD	INDIA

Appln Info Contents Petition Info Atty/Agent Info Continuity/Reexam	Foreign [
Search Another: Application# Search or Patent# Sea	arch
PCT / Search or PG PUBS #	Search
Attorney Docket # Search	
Bar Code # Search	

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         NOV 10
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         NOV 10
NEWS
                 8.01c now available
                 CA/CAplus to MARPAT accession number crossover limit increased
         NOV 20
NEWS
                 to 50,000
                 CAS REGISTRY updated with new ambiguity codes
         DEC 01
NEWS
                 CAS REGISTRY chemical nomenclature enhanced
NEWS 10
         DEC 11
                 WPIDS/WPINDEX/WPIX manual codes updated
         DEC 14
NEWS 11
                 GBFULL and FRFULL enhanced with IPC 8 features and
         DEC 14
NEWS 12
                 functionality
                 CA/CAplus pre-1967 chemical substance index entries enhanced
         DEC 18
NEWS 13
                 with preparation role
                 CA/CAplus patent kind codes updated
NEWS 14
         DEC 18
                 MARPAT to CA/CAplus accession number crossover limit increased
         DEC 18
NEWS 15
                 to 50,000
                 MEDLINE updated in preparation for 2007 reload
         DEC 18
NEWS 16
                 CA/CAplus enhanced with more pre-1907 records
         DEC 27
NEWS 17
                 CHEMLIST enhanced with New Zealand Inventory of Chemicals
         JAN 08
NEWS 18
                 CA/CAplus Company Name Thesaurus enhanced and reloaded
         JAN 16
NEWS 19
                 IPC version 2007.01 thesaurus available on STN
         JAN 16
NEWS 20
                 WPIDS/WPINDEX/WPIX enhanced with IPC 8 reclassification data
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         JAN 16
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         JAN 22
NEWS 22
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         JAN 22
                 PHAR reloaded with new search and display fields
         JAN 29
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                 CAS Registry Number crossover limit increased to 300,000 in
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                  multiple databases
                  CASREACT coverage to be extended
NEWS 26
         FEB 13
                  PATDPASPC enhanced with Drug Approval numbers
NEWS 27
          Feb 15
                  RUSSIAPAT enhanced with pre-1994 records
         Feb 15
NEWS 28
                  KOREAPAT enhanced with IPC 8 features and functionality
NEWS 29 Feb 23
              NOVEMBER 10 CURRENT WINDOWS VERSION IS V8.01c, CURRENT
 NEWS EXPRESS
               MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
               AND CURRENT DISCOVER FILE IS DATED 25 SEPTEMBER 2006.
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chain nodes:
7 20 21 22 23 24 25 26 27 28

ring nodes:
1 2 3 4 5 6 8 9 10 11 12 13 14 15 16 17 18 19

chain bonds:
6-7 7-8 7-14 11-28 17-20 20-21 21-22 22-23 23-24 24-25 24-26 25-27

ring bonds:
1-2 1-6 2-3 3-4 4-5 5-6 8-9 8-13 9-10 10-11 11-12 12-13 14-15 14-19
15-16 16-17 17-18 18-19

exact/norm bonds:
7-14 14-15 14-19 15-16 16-17 17-18 17-20 18-19 21-22 22-23

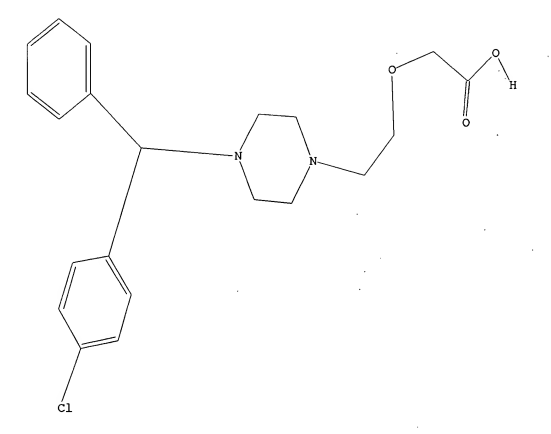
exact bonds:
6-7 7-8 11-28 20-21 23-24 25-27

normalized bonds:
1-2 1-6 2-3 3-4 4-5 5-6 8-9 8-13 9-10 10-11 11-12 12-13 24-25 24-26

Match level:
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:Atom 20:CLASS 21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:CLASS 27:CLASS 28:CLASS

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18 ITERATIONS

4 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

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PROJECTED ITERATIONS:

106 TO 614

PROJECTED ANSWERS:

4 TO

L2

4 SEA SSS SAM L1

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FULL SEARCH INITIATED 18:42:18 FILE 'REGISTRY.'

FULL SCREEN SEARCH COMPLETED - 374 TO ITERATE

100.0% PROCESSED

374 ITERATIONS

41 ANSWERS

SEARCH TIME: 00.00.01

L3

41 SEA SSS FUL L1

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SINCE FILE TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

172.10

172.31

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=> s 13

L4 1053 L3

=> S (CRYSTALLINE)

L5 77110 (CRYSTALLINE)

=> S (L4)

L6 1053 (L4)

=> S L6 AND (CRYSTALLINE)

77110 CRYSTALLINE

L7 5 L6 AND (CRYSTALLINE)

=> S L6 AND (MONOHYDROCHLORIDE)

3690 MONOHYDROCHLORIDE

L8 4 L6 AND (MONOHYDROCHLORIDE)

=> S L6 AND (XRAY)

4684 XRAY

L9 0 L6 AND (XRAY)

=> S L6 AND (X-RAY)

1574267 X

1063009 RAY

821830 X-RAY

(X (W) RAY)

L10 5 L6 AND (X-RAY)

=> S L6 AND (DIFFRACTION)

446102 DIFFRACTION

L11 4 L6 AND (DIFFRACTION)

=> s 17 or 18 or 110 or 111

L12 10 L7 OR L8 OR L10 OR L11

=> s 112 not (2006/so or 2005/so)

795084 2006/SO

867953 2005/so

L13 10 L12 NOT (2006/SO OR 2005/SO)

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L13 ANSWER 1 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2006:1329533 CAPLUS

DOCUMENT NUMBER:

146:87555

TITLE:

Cetirizine hydrochloride masticatory tablet and its

preparation

INVENTOR(S):

Gu, Xuchu; Zhong, Xuebin

PATENT ASSIGNEE(S):

Nanjing Golden Eagle Medicinery Technology Development

Co., Ltd., Peop. Rep. China

SOURCE:

Faming Zhuanli Shenqing Gongkai Shuomingshu, 6pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent

LANGUAGE:

Chinese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1875970	A	20061213	CN 2005-10040439 CN 2005-10040439	20050608 20050608

83881-52-1, Cetirizine hydrochloride

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (Cetirizine hydrochloride masticatory tablet and its preparation)

83881-52-1 CAPLUS RN

Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-, CN dihydrochloride (9CI) (CA INDEX NAME)

2 HCl

The title masticatory tablet is composed of cetirizine hydrochloride 1, AB crystalline cellulose 10-50, sucrose 10-50, β -cyclodextrin 1-10, sodium carboxymethyl starch 1-10, sodium saccharin 0.2-5, magnesium stearate 0.1-10, micropowder silica gel 0.1-5 part, and water proper quantity. preparation method comprises pulverizing, sieving by 80 mesh sieve, mixing cetirizine hydrochloride, sodium saccharin, β-cyclodextrin, sucrose powder, and crystalline cellulose with distilled water to obtain soft material, sieving by 40 mesh sieve, prilling, drying at 60° for 2 h, sieving by 30 mesh sieve, adding sodium carboxymethyl starch, magnesium stearate, and micropowder silica gel, stirring, and pressing. The invention can be used for treating seasonal or perennial allergic rhinitis, and urticaria and cutaneous pruritus caused by allergen.

L13 ANSWER 2 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2006:381409 CAPLUS

DOCUMENT NUMBER:

144:432829

TITLE:

Preparation of 2,6-substituted-4-monosubstituted amino-pyrimidines as prostaglandin D2 receptor

antagonists

INVENTOR(S):

Lim, Sungtaek; Harris, Keith John; Stefany, David; Gardner, Charles J.; Cao, Bin; Boffey, Ray; Gillespy, Timothy A.; Aguiar, Joacy C.; Hunt, Hazel J.; Dechaux,

Elsa A.

PATENT ASSIGNEE(S): SOURCE:

Aventis Pharmaceuticals Inc., USA

PCT Int. Appl., 272 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

•	PATE	I TN	10.			KINI		DATE		1	APPL	CAT:	ION I	10. 					
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The invention is directed to the preparation of aminopyrimidines I [Cy1 = AB (un) substituted cycloalkyl, heterocyclyl, hetero/aryl, etc.; Cy2 = (un) substituted cycloalkenyl, heterocyclenyl, hetero/aryl, etc.; L1 = cyclo/alkylene, CH2-haloalkylene; or L1Cy2 = arylcycloalkyl, cycloalkylaryl; R1 = alkylthio, NH2 and derivs., alkoxy; L2 = a bond, O, CH2O; provided that when R1 = OMe, L1 = CH2CH2, L2 = a bond, and Cy2 = 2,4-dichlorophenyl, then Cyl is not 1-methyl-2-ethyloxycarbonylindol-5yl], and their N-oxides, ester prodrugs, and their pharmaceutically acceptable salts, hydrates and solvates, and their use as prostaglandin D2 (PGD2) receptor antagonists in pharmaceutical compns. comprising a pharmaceutically effective amount of one or more compds. I in admixt. with a pharmaceutically acceptable carrier, and to a method of treating a patient suffering from a PGD2-mediated disorder. E.g., a 4-step synthesis, starting from from 3-fluoro-4-methoxybenzaldehyde, was given for pyrimidine II. Selected I produced 50% inhibition in the SPA cAMP assay in human LS174T cells expressing the endogenous DP receptor at concns. within the range of about 0.1 to about 30 nM. I are useful for treating allergic disease (such as allergic rhinitis, allergic conjunctivitis, atopic dermatitis, bronchial asthma and food allergy), systemic mastocytosis, disorders accompanied by systemic mast cell activation, anaphylaxis shock, bronchoconstriction, bronchitis, urticaria, eczema, diseases accompanied by itch, diseases (such as cataract, retinal detachment, inflammation, infection and sleeping disorders) which are generated secondarily as a result of behavior accompanied by itch (such as scratching and beating), chronic obstructive pulmonary diseases, ischemic reperfusion injury, cerebrovascular accident, chronic rheumatoid arthritis, pleurisy, ulcerative colitis (no data):

L13 ANSWER 3 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

2005:523256 CAPLUS ACCESSION NUMBER:

143:65406 DOCUMENT NUMBER:

Multiparticulate crystalline drug TITLE:

II

compositions containing a Poloxamer and a glyceride Appel, Leah Elizabeth; Crew, Marshall David; Friesen, INVENTOR(S): Dwayne Thomas; Herbig, Scott M.; Lo, Julian Belknap; Lyon, David Keith; McCray, Scott Bladwin; Ray,

Roderick Jack; West, James Blair

PATENT ASSIGNEE(S):

Pfizer Products Inc., USA

SOURCE:

PCT Int. Appl., 46 pp.

DOCUMENT TYPE:

Patent

CODEN: PIXXD2

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT NO.				KIN	D	DATE			APPL	ICAT:	ION 1	NO.	_	DATE				
		2005 2005									WO 2	004-	IB38	08		20041122			
											BB,	BG.	BR.	BW.	BY,	BZ.	CA,	CH.	
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A multiparticulate for controlled release of a crystalline drug comprises a AB glyceride having at least one alkylate substituent of at least 16 carbon atoms, and a Poloxamer, wherein at least 70 weight% of the drug in the multiparticulate is crystalline Thus, azithromycin-containing

multiparticulates were prepared via a melt-congeal process from a mixture containing azithromycin/Compritol 888 ATO/Pluronic (50:40:10) forming a preblend and extrusion of the preblend at a feed rate of 130 g/min. More than 90 weight% of the azithromycin in the multiparticulates was crystalline dihydrate. release rate of azithromycin from the multiparticulates was 32, 67, 90,

99, 99, and 100% in 5, 15, 30, 60, 120, and 180 min, resp.

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2007 ACS on STN L13 ANSWER 4 OF 10

6

ACCESSION NUMBER:

2005:310383 CAPLUS

DOCUMENT NUMBER:

143:159082

TITLE:

Preparation of inclusion complex of cetirizine- β -

cyclodextrin

AUTHOR(S):

Zhao, Liping; Cao, Deying; Feng, Xiangping; Yuan,

CORPORATE SOURCE:

Pharmacy College, Hebei Medical University,

Shijiazhuang, 050017, Peop. Rep. China Huaxi Yaoxue Zazhi (2004), 19(1), 30-32

CODEN: HYZAE2; ISSN: 1006-0103

PUBLISHER: Huaxi Yike Daxue Yaoxueyuan

DOCUMENT TYPE: Journal LANGUAGE: Chinese

IT 83881-52-1, Cetirizine hydrochloride

RL: BSU (Biological study, unclassified); BIOL (Biological study) (preparation of inclusion complex of cetirizine- β - cyclodextrin)

RN 83881-52-1 CAPLUS

SOURCE:

CN Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-, dihydrochloride (9CI) (CA INDEX NAME)

●2 HCl

AB The inclusion compound of cetirizine- β -cyclodextrin was prepared and its properties were studied. The inclusion compound was prepared by saturated solution

method, and it was proved by the changes of physics properties before and after inclusion. The average inclusion rate of the inclusion compound was 85.1%. The UV spectra, solubility, taste, stability, X-ray diffractometry and IR spectra showed that the inclusion compound became a new complex. The inclusion compound of cetirizine- β -cyclodextrin could cover the no good taste of cetirizine.

L13 ANSWER 5 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:238545 CAPLUS

DOCUMENT NUMBER: 142:291446

TITLE: Methods and kits for monitoring resistance to

therapeutic agents Cantor, Thomas L.

INVENTOR(S): Cantor,

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 24 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2005059023 PRIORITY APPLN. INFO.:	A1 :	20050317	05 2005 001200	20030916 20030916

IT 83881-51-0, Cetirizine 83881-52-1, ZYRTEC

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (methods and kits for monitoring resistance to therapeutic agents)

RN 83881-51-0 CAPLUS

CN Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy](9CI) (CA INDEX NAME)

83881-52-1 CAPLUS RN

Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-, CN dihydrochloride (9CI) (CA INDEX NAME)

2 HCl

The invention relates to novel methods and kits for monitoring the AB therapeutic inactivating capacity of a subject. The invention further relates to methods and kits for determining and/or monitoring a therapeutic protocol for a subject afflicted with auto-antibodies specific for a natural substance, wherein these auto antibodies develop as a result of therapeutic administration of the natural substance or an analog thereof. These methods and kits can be used, for example, to initiate, terminate, or adjust the level of administration of any of a variety of therapeutic agents.

L13 ANSWER 6 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2005:78236 CAPLUS

DOCUMENT NUMBER:

142:162672

TITLE:

Crystalline cetirizine

monohydrochloride

INVENTOR(S):

Reddy, Manne Satyanarayana; Rajan, Srinivasan

Thirumalai; Rao, Uppala Venkata Bhaskara; Reddy, Konda

Srinivasa

PATENT ASSIGNEE(S):

Reddy's Laboratories Limited, India; Reddy's

Laboratories, Inc.

SOURCE:

U.S. Pat. Appl. Publ., 11 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND .	DATE	APPLICATION NO.	DATE
us 2005020608	A1	20050127	US 2004-809192	20040325
IN 2003MA00252	Α	20050304	IN 2003-MA252	20030325
PRIORITY APPLN. INFO.:			IN 2003-MA252 A	2.0030325

798544-25-9P TT ·

RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation of crystalline cetirizine monohydrochloride for oral dosage forms)

RN 798544-25-9 CAPLUS

Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-, CN monohydrochloride (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{C1} & \text{Ph} & \text{CH}_2\text{-}\text{CH}_2\text{-}\text{O}\text{-}\text{CH}_2\text{-}\text{CO}_2\text{H} \\ & \text{CH} & \text{N} & \text{N} \end{array}$$

HCl

83881-51-0P, Cetirizine IT RL: RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation of crystalline cetirizine monohydrochloride for oral dosage forms)

83881-51-0 CAPLUS RN

Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-CN (CA INDEX NAME) (9CI)

83881-52-1P, Cetirizine dihydrochloride IT RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of crystalline cetirizine monohydrochloride for oral dosage forms)

83881-52-1 CAPLUS RN

Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-, CN dihydrochloride (9CI) (CA INDEX NAME)

2 HCl

A novel crystalline form of cetirizine monohydrochloride and processes for making the crystalline form as well as compns., pharmaceutical compns., and methods utilizing the crystalline form are described. A process for preparation of a crystalline form of cetirizine monohydrochloride, comprises (1) providing a solid residue of crude cetirizine. monohydrochloride; (2) contacting the crude residue with a ketone solvent to cause separation of a solid mass; and (3) isolating the solid mass thereby obtaining the crystalline form of cetirizine monohydrochloride Tablets for the treatment of allergic syndromes were formulated containing crystalline cetirizine monohydrochloride 10, CaCO3 500, PVP 17, Avicel 15, mannitol 400, maltodextrin 15, aspartame 3, and aroma 20 mg each.

L13 ANSWER 7 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

2004:1037084 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 142:6558 Preparation of 2-[2-[4-[(4-chlorophenyl)phenylmethyl]-TITLE: 1-piperazinyl]ethoxy]acetic acid monohydrochloride as antiallergic compound Singh, Shiva Prasad; Mukarram, Siddiqui Mohammed INVENTOR(S): Jaweed; Merwade, Aravind Yekanathsa; Khan, Anjum Reyaz Wockhardt Limited, India PATENT ASSIGNEE(S): PCT Int. Appl., 31 pp. SOURCE: CODEN: PIXXD2 DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: DATE APPLICATION NO. KIND DATE PATENT NO. _____ 20030521 20041202 WO 2003-IB1947 WO 2004103982 A1 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG AU 2003-228011 20030521 20041213 A1 AU 2003228011 EP 2003-725479 20030521 A1 20060301 EP 1628964 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

IE, SI, FI, RO, CY, TR, BG, CZ, EE, HU, SK US 2006-554696 20060223 20061116 US 2006258684 A1 A 20030521 WO 2003-IB1947 PRIORITY APPLN. INFO .: CASREACT 142:6558 OTHER SOURCE(S): 798544-25-9P RL: IMF (Industrial manufacture); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of 2-[2-[4-[(4-chlorophenyl)phenylmethyl]-1piperazinyl]ethoxy]acetic acid monohydrochloride as

antiallergic compound) RN 798544-25-9 CAPLUS

CN Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-, monohydrochloride (9CI) (CA INDEX NAME)

• HCl

IT 83881-52-1

RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of 2-[2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]acetic acid monohydrochloride as antiallergic compound)

RN 83881-52-1 CAPLUS

Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-, CN dihydrochloride (9CI) (CA INDEX NAME)

2 HCl

83881-51-0P, Cetirizine IT

RL: RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation of 2-[2-[4-[(4-chlorophenyl)phenylmethyl]-1piperazinyl]ethoxy]acetic acid monohydrochloride as antiallergic compound)

83881-51-0 CAPLUS RN

Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-CN (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{C1} & \text{Ph} & \text{CH}_2\text{--}\text{CH}_2\text{--}\text{O}\text{--}\text{CH}_2\text{--}\text{CO}_2\text{H} \\ \hline & \text{CH} & \text{N} \end{array}$$

The title compound (I) is prepared by reaction of 4-chlorobenzhydrylpiperazine AΒ with 2-chloroethanol followed by reaction with sodium chloroacetate and salt formation. I was characterized by DSC, NMR, X-ray powder diffraction, m.p., elemental anal., and HPLC.

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 8 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

3

ACCESSION NUMBER:

2004:493694 CAPLUS

DOCUMENT NUMBER:

141:54360

TITLE:

Polymorphic crystalline forms of

dihydrochloride salts of cetirizine and processes for

their preparation

INVENTOR(S):

Reddy, Manne Satyanarayana; Srinivasan, Thirumalai

Rajan; Uppala, Venkata Bhaskara Rao; Vaddadi, Pattabhi Ramayya; Joga, Rajender

PATENT ASSIGNEE(S):

Reddy's Laboratories Limited, India; Reddy's

Laboratories, Inc.

SOURCE:

PCT Int. Appl., 37 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004050647 WO 2004050647	A2 A3	20040617 20040902	WO 2003-US38494	20031204
WO 2004050647	A8	20050303 , AU, AZ, BA	, BB, BG, BR, BW, BY,	BZ, CA, CH,

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CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
             GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
             LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO,
                         PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ,
             NZ, OM, PG,
                         TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
             TM, TN,
                     TR,
                         KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
         RW: BW, GH,
                     GM,
                         MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE,
             BY, KG,
                     KZ,
                            GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK,
                FI,
                    FR,
                         GB,
                             CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                    ВJ,
                         CF,
             TR, BF,
     IN 2002MA00908
                          Α
                                20050304
                                             IN 2002-MA908
                                                                     20021204
                                20040617
                                             CA 2003-2488114
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     CA 2488114
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    AU 2003297640
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                                20040623
                                             AU 2003-297640
                                                                     20031204
                                             US 2003-729856
                                                                     20031204
     US 2004186112
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                                20040923
                          Α
                                20051102
                                             CN 2003-80100543
                                                                     20031204
     CN 1692105
                                             IN 2002-MA908
                                                                 Α
                                                                     20021204
PRIORITY APPLN. INFO.:
                                             WO 2003-US38494
                                                                 W
                                                                     20031204
     130018-87-0P 163837-48-7P
IT
     RL: PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
        (polymorphic crystalline forms of dihydrochloride salts of cetirizine and
        processes for their preparation)
     130018-87-0 CAPLUS
RN
     Acetic acid, [2-[4-[(R)-(4-chlorophenyl)phenylmethyl]-1-
CN
```

Absolute stereochemistry. Rotation (+).

●2 HCl

RN 163837-48-7 CAPLUS

CN Acetic acid, [2-[4-[(S)-(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-, dihydrochloride (9CI) (CA INDEX NAME)

piperazinyl]ethoxy]-, dihydrochloride (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

●2 HCl

IT 130018-76-7P, Dextrocetirizine 130018-77-8P,
 Levocetirizine
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
 (Reactant or reagent)

(polymorphic crystalline forms of dihydrochloride salts of cetirizine and processes for their preparation from)

RN 130018-76-7 CAPLUS

CN Acetic acid, [2-[4-[(S)-(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 130018-77-8 CAPLUS

CN Acetic acid, [2-[4-[(R)-(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

AB Crystalline polymorphic forms of the levorotatory and dextrorotatory cetirizine dihydrochloride salts are prepared by dissolving the salts in an a ketone-containing solvent (e.g., aqueous acetone), cooling the solution, and collecting the crystalline precipitate

L13 ANSWER 9 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2003:991495 CAPLUS

DOCUMENT NUMBER:

140:47519

TITLE:

Process for the preparation of an amorphous form of

[2-[4-[(4-chlorophenyl)phenylmethyl]-1-

piperazinyl]ethoxy]acetic acid dihydrochloride

(cetirizine dihydrochloride)

INVENTOR(S):

Reddy, Manne Satyanarayana; Rajan, Srinivasan Thirumalai; Shankar, Ranga Ravi; Vardhan, Sunkara

Vishnu

PATENT ASSIGNEE(S):

Dr.Reddy's Laboratories Ltd., India; Dr.Reddy's

Laboratories, Inc.

SOURCE:

PCT Int. Appl., 21 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

2

PATENT INFORMATION:

PATENT NO. KIND					D :	DATE	•	APPLICATION NO.							DATE		
WO 2003104212 A1						20031218 WO 2003-US17600							20030604				
W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
	co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,	
	LS.	LT.	LU.	LV,	MA.	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	NΖ,	OM,	

PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG AU 2003-238883 20030604 20031222 AU 2003238883 A1 IN 2002-MA425 20020605 PRIORITY APPLN. INFO .: 20030604 WO 2003-US17600

IT 83881-51-0P, Cetirizine

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(in a process for the preparation of an amorphous form of

[2-[4-[(4-chlorophenyl)phenylmethyl]-1- piperazinyl]ethoxy]acetic acid dihydrochloride (cetirizine dihydrochloride))

RN 83881-51-0 CAPLUS

CN Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-(9CI) (CA INDEX NAME)

IT 83881-52-1P, Cetirizine dihydrochloride
 RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP
 (Physical process); SPN (Synthetic preparation); PREP (Preparation); PROC
 (Process)

(process for the preparation of an amorphous form of [2-[4-[(4-chlorophenyl)phenylmethyl]-1- piperazinyl]ethoxy]acetic acid dihydrochloride (cetirizine dihydrochloride))

RN 83881-52-1 CAPLUS

CN Acetic acid, [2-[4-[(4-chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]-, dihydrochloride (9CI) (CA INDEX NAME)

●2 HCl

AB A novel, amorphous form of [2-[4-[(4-Chlorophenyl)phenylmethyl]-1-piperazinyl]ethoxy]acetic acid dihydrochloride, suitable for pharmaceutical formulations, is prepared and X-ray diffraction patterns for it are presented.

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 10 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2003:991494 CAPLUS

DOCUMENT NUMBER:

140:42205

TITLE:

Preparation of crystalline

[2-[4-[(4-chlorophenyl)phenylmethyl]-1-

piperazinyl]ethoxy]acetic acid dihydrochloride

(cetirizine dihydrochloride)

INVENTOR(S):

Reddy, Manne Satyanarayana; Rajan, Srinivasan

Thirumalai; Shankar, Ranga Ravi; Vardhan, Sunkara

Vishnu

PATENT ASSIGNEE(S): Reddy's Laboratories Limited, India; Reddy's

Laboratories, Inc.

SOURCE: PCT Int. Appl., 22 pp.

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE:

: Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT				KINI		DATE		-	APPL:							
	WO 200	31042	11		A2		2003	1218	1	WO 2					20030604		
		31042	AG,	ът	AM A	ידי ע	2004	Δ7.	RΔ	BB.	BG.	BR.	BY.	BZ.	CA.	CH.	CN,
	W:	AL,	CR,	AL,	Au,	VI,	nv,	DM	D71,	EC,	EE.	ES.	FT.	GB.	GD.	GE.	GH.
		co,	HR,	CU,	CZ	DE,	DA,	DM,	TD,	EC,	KC	KD,	KD	K7.	T.C	T.K.	LR.
		GM,	HR,	HU,	ID,	TL,	TN,	15,	UE,	MAI,	MG,	MY.	M7	MT	NO.	NZ	OM.
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		TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW	73.6	77.17	7\M	7.7	RV
	RV	: GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	ou,	ΔM,	ΔW,	AM,	RU,	EC.
		KG,	KZ,	MD,	RU,	ТJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	ee,	ωD TO'
		FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PT,	KO,	DE,	SI,	DV,	Tr,
		BF,	BJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MK,	ΝĿ,	211,	10,	CO4
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										WO 2	003-	USI7	6/2		w 2	0030	604
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	piperazinyl]ethoxy]acetic acid dihydrochloride (cetirizine																
		nydrod															
RN	83881	-52-1	CAP	TJUS				•									
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		1.1															

$$\begin{array}{c|c} \text{C1} & \text{Ph} & \text{CH}_2\text{--}\text{CH}_2\text{--}\text{O}\text{--}\text{CH}_2\text{--}\text{CO}_2\text{H} \\ \hline \\ \text{CH} & \text{N} & \end{array}$$

dihydrochloride (9CI) (CA INDEX NAME)

•2 HCl

$$\begin{array}{c|c} \text{Cl} & \text{Ph} & \text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CO}_2\text{H} \\ \hline & \text{CH}-\text{N} & \text{N} \end{array}$$

AB A crystalline form of cetirizine dihydrochloride (I), prepared by the salification of cetirizine with isopropanolic hydrogen chloride, having a defined X-ray diffraction pattern is presented, and pharmaceutical compns. containing I are presented.

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---Logging off of STN---

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Executing the logoff script...

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COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	75.02	247.33
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	-7.80	-7.80

STN INTERNATIONAL LOGOFF AT 18:47:34 ON 23 FEB 2007